Research Paper

Assessment of Groundwater Quality in the Jajrood River Basin, Tehran, Iran: A Coupled Physicochemical and Hydrogeochemical Study

Maryam Givi1, Mahsa Jahangiri-Rad2*, Hamidreza Tashauoei1

1. Department of Environmental Health Engineering, Faculty of Public Health and Biomedical Engineering, Tehran Medical Sciences, Islamic Azad University, Tehran, Iran
2. Water Purification Research Center, Tehran Medical Sciences, Islamic Azad University, Tehran, Iran.

* Corresponding Author:
Mahsa Jahangiri-Rad, PhD.
Address: Water Purification Research Center, Tehran Medical Sciences, Islamic Azad University, Tehran, Iran.
Phone:+98 (912) 2834504
E-mail: mahsajahangiri_64@yahoo.com; m.jahangiri@iautmu.ac.ir

Abstract

Background: The physicochemical composition of groundwater is affected by the quantity and quality of surrounding aquifers which are in turn recharging from adjacent river waters.

Methods: In the present study, 20 surface and 16 groundwater samples were collected in pre- and post-monsoon season from the Jajrood River basin, Tehran, Iran. The samples were analyzed for 18 physicochemical water quality characteristics to assess the river and groundwater qualities. Hydrogeochemical analyses of groundwater samples were also performed to determine the Water Quality Index (WQI) for drinking and evaluate factors governing the water quality characteristic in the study area. Accordingly, the Piper diagram and Gibbs and Chadha plots were drawn to assess seasonal variations in hydrochemical facies and processes in the basin. Subsurface soil samples were also examined with respect to the structure, elemental composition, and multi-elemental trace analysis.

Results: Results showed the abundance of major ions in the order of Ca²⁺ >Na⁺>Mg²⁺>K⁺ for cations and HCO⁻₃ >SO₄²⁻ >Cl⁻ >NO₃⁻ >F⁻ for anions. In general, all drinking groundwater samples met WHO permissible limits except for Chemical Oxygen Demand (COD) and HCO⁻₃. Moreover, the water is categorized as Ca-Mg-HCO₃ type. Subsurface soil analyses demonstrated quartz and calcium carbonate as the main phases of soil structure, suggesting the enrichment of groundwater with temporary hardness.

Conclusion: Overall, the groundwater quality was suitable for drinking and agricultural activities.
1. Introduction

Groundwater virtually accounts for half of all drinking water all over the world [1]. The quality of water through its biological content and chemical composition affects public health and agricultural production [2]. In recent decades, the quality of water bodies has decreased either by anthropogenic or natural processes, which subsequently has deteriorated its use for drinking, agriculture, and other purposes [3, 4]. The general hydrogeochemical characteristics of water regulate its use for intended purposes. Water chemistry is mainly influenced by the interaction of water with lithologic media through which it flows [5]. The chemical composition of groundwater, therefore, depends on the quality of surface water in the recharge zone and also upon various hydrogeochemical processes that happen naturally in the subsurface [6]. Seasonal variations in groundwater chemistry are also contributed to these processes. Moreover, soils have important direct and indirect impacts on water quality. A natural phenomenon, like topographic position and the mineral composition of the underlying geology, is responsible for the primary physical and geochemical conditions in groundwater that account for physical properties, such as pH, temperature, electrical conductivity, alkalinity, and chemical concentrations of dissolved solids [7].

The chief ion profile of groundwater can provide accurate information to ascertain solute sources and delineate groundwater evolution [8, 9]. Heretofore, hydrochemical facies have been widely considered in the chemical evaluation of groundwater and surface water. In this regard, Piper [10] and Durov [11] conducted the first investigation. After that, Chadha [12] developed a simple modified Piper diagram. Many researchers have extensively used these diagrams to scrutinize the driving factors of water chemistry [13-15]. Several authors have used the Water Quality Index (WQI) to assess the water chemistry status in the river and groundwater for human consumption [16, 17].

WQI provides a standardized method to compare among diverse water sources based upon water quality parameters, such as common anions and cations, pH, electrical conductivity, turbidity, fecal coliforms, etc. Based on this index, the water quality is categorized as excellent, good, medium, bad, and very bad [18]. Several studies have been conducted in Iran to understand the causes and effects of aquifers-related issues besides their current groundwater characteristics. These studies suggest that various processes influence groundwater chemistry and its impact on different beneficial uses (drinking, industrial, agricultural) [19-21]. A large and growing body of literature has investigated the surface water and groundwater interaction using hydrogeology and hydrochemical assessment. Piper, Gibbs, and Chadha diagrams have been reported as robust tools for the hydrogeochemical evaluation since the former can classify the groundwater and the latter estimates the origin of ions [22-24].

In many rural areas of Iran, wells provide a reliable and ample water supply for domestic uses. Several dug wells have recently been excavated in Pardis City, along the Jajrood river, to provide water for drinking and irrigation purposes. Drilling other wells in this area entails checking the water quality of present dug wells that are supplied by the river. The present study investigated the quality of the Jajrood river water and its effect on the physical and chemical properties of shallow wells in the study area.

An initial objective of this research was to analyze the hydrogeochemical characteristics of the shallow groundwater and their evolution using descriptive statistics and the Gibbs diagram. Furthermore, water types and the corresponding controlling factors of groundwater characteristics were assessed using the Piper, and Chadha approaches for the pre- and post-monsoon samples. Since the soil analyses through which the water passes provide reliable information about the chemical properties of water, we examined the structure and elements of soil samples, too. Finally, the role of subsurface soil in the chemistry of groundwater constituents was investigated.

Details of the study site

Study area

The study area is part of the Jajrood catchment in Tehran Province based on urban political divisions (Figure 1). The geographical location of the basin is 51°5’ to 51°24’ east longitude and 35°46’ to 36°3’ north latitude. The river originates from the heights of the Alborz mountain range in the north of Tehran. The river is located 30 km northeast of Tehran City, the capital of Iran, with a length of 40 km and 710 km² catchment area, which flows from northwest to southeast. On its way, the river supplies drinking and agricultural water for many villages, and after joining the Garmabdar branch and Ruteh river to the southeast, it enters the Latyan dam and ends at Namak (Salt) Lake. The average annual rainfall of the study area is recorded as almost 600 mm [25]. The decrease in temperature is observed from south to north of the research area. The average annual temperature is less than 5 °C in the highlands and 21 °C in the south-
ern parts. The relative humidity of the river basin varies between 18% to 45%. Pardis city, consisting of Bumhan and Jajrood districts, is in this catchment area. This city has three districts of Bumhan, Markazi, and Jajrood. The Jajrood district includes Jajrood village and Saeedabad district, where all the samples in the present study were taken.

Geology and hydrology

The general terrain of the study area consists mainly of some faults and folds to the south [25]. The soil of the area is mostly alluvial sediments and includes sand and gravel. The basin of the Jajrood river is in an area with a maximum altitude of 4000 m above sea level. The area can be divided into two parts in terms of elevation: 1) the mountainous area, which forms a smaller part of the Jajrood river, encompasses the continuation of the Central Alborz mountain range along with hills part of which stretches into a forest, and 2) the plain area, which is the large part of the river's catchment area. The latter is not very high, and only in some areas, little vegetation can be observed.

2. Materials and Methods

Field survey and chemical analysis

Twenty surface water and 16 groundwater samples were taken in pre- and post-monsoon seasons from 5 points spread over the Jajrood river basin and 4 wells recharging from the river (Figure 1). Sampling was done twice from each point, and the average results were reported. For chemical analysis, water samples were collected in polyethylene bottles. The sampling bottles were soaked with HCl solution then washed thoroughly twice before sampling. pH and Electrical Conductivity (EC) were measured in situ by portable digital meters. For cation analysis, the samples were filtered through a 0.45 μm cellulose acetate membrane, and nitric acid was added to the bottles to reach a pH below 2. All samples were refrigerated and analyzed within 12 hours. The sampling was followed based on standard protocols prescribed by American Public Health Association (APHA) [26]. The measurements were repeated twice, and the arithmetic average of the two measurement results was reported.

Bacteriological analysis was conducted in the same river and groundwater samples of the study area in pre- and post-monsoon seasons. The sodium thiosulfate rinsed bottles which were further sterilized in an incubator, were used for microbial sample collection. The 9-tube fermentation technique was used, and the total number of coliforms was determined as the most probable number per 100 mL (MPN/100 mL). The analytical methods for each parameter are summarized in Table 1. Piper pilot (Piper, 1944) and Chadha’s (1999) approach were applied to determine groundwater types of the study area. Gibbs (1970) diagram was used to establish the relationship of water composition and aquifer lithological characteristics [27]. All diagrams were prepared using Microsoft Excel v. 2013.

Water Quality Indexes

WQI expresses as a rating that reflects the mixed influence of various water quality parameters [28]. To calculate this index, we assigned each water quality parameter an assorted weight (wi) on a scale of 1 to 5 based on the importance of their health effects and the significance in drinking water quality (Table 2). High weight values are assigned to parameters that, if exceeded by the standard limits, have severe health effects and make the water unusable for drinking purposes [29]. The relative weight (Wi) is computed from the Equation 1 [30]:

\[ W_i = \frac{w_i}{\sum w_i} \]

where, \( W_i \) represents weight, \( w_i \) is the weight of each parameter and shows the sum of all parameters included in the WQI calculation.

Then, a quality rating (qi) for each parameter was obtained by dividing its concentration (Ci) by its limit values (Si) given in Table 2 based on Equation 2 [17].

\[ q_i = \frac{C_i}{S_i} \times 100 \]

To calculate WQI, the SIi value was calculated according to Equation 3:

\[ S_{ii} = W_i \times q_i \]

Ultimately, WQI was computed by Equation 4 [17].

\[ WQI = \sum_{i=1}^{n} S_{ii} \]

Sodium percentage (Na%), sodium absorption ratio (SAR), and residual sodium carbonate (RSC) were computed to investigate the groundwater suitability of Gw2 for irrigation purposes. The concentration of groundwater samples in the following equations is presented in meq/L. SAR was obtained from the Equation 5 [31].

\[ \text{SAR} = \frac{Na}{Ca+Mg} \]

Na% was computed using Equation 6 [32]:

\[ \text{Na%} = \frac{\text{Na}}{\text{Ca}+\text{Mg}} \times 100 \]
Table 1. Sampling analysis techniques

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Methods</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cations</td>
<td>Induced Coupled Plasma (ICP)</td>
</tr>
<tr>
<td>Anions</td>
<td>Ion Chromatograph (IC)</td>
</tr>
<tr>
<td>Alkalinity</td>
<td>Acid titration</td>
</tr>
<tr>
<td>Total Hardness (TH)</td>
<td>EDTA titration</td>
</tr>
<tr>
<td>Total Dissolved Solids (TDS)</td>
<td>Mathematical calculations (using anions and cations concentration)</td>
</tr>
<tr>
<td>pH</td>
<td>Electrometric</td>
</tr>
<tr>
<td>Turbidity</td>
<td>Nephelometry</td>
</tr>
<tr>
<td>Electrical conductivity (EC)</td>
<td>Conductivity meter</td>
</tr>
<tr>
<td>Zn</td>
<td>Polarography</td>
</tr>
<tr>
<td>Chemical Oxygen Demand (COD)</td>
<td>Spectrometry</td>
</tr>
<tr>
<td>Coliforms</td>
<td>9- tubes (Multi-pipe fermentation)</td>
</tr>
</tbody>
</table>

Table 2. Descriptive Statistics of parameters and the values used for Water Quality Index (WQI) calculation

<table>
<thead>
<tr>
<th>Parameters</th>
<th>WHO (2011)</th>
<th>Weight (wi)</th>
<th>Relative Weight (Wi)</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH</td>
<td>6.5-8.5</td>
<td>4</td>
<td>0.085</td>
</tr>
<tr>
<td>EC</td>
<td>1000</td>
<td>4</td>
<td>0.085</td>
</tr>
<tr>
<td>TDS</td>
<td>500</td>
<td>5</td>
<td>0.106</td>
</tr>
<tr>
<td>TH</td>
<td>200</td>
<td>2</td>
<td>0.042</td>
</tr>
<tr>
<td>F</td>
<td>1.5</td>
<td>5</td>
<td>0.106</td>
</tr>
<tr>
<td>Cl</td>
<td>250</td>
<td>3</td>
<td>0.063</td>
</tr>
<tr>
<td>SO\textsubscript{4}</td>
<td>250</td>
<td>4</td>
<td>0.085</td>
</tr>
<tr>
<td>HCO\textsubscript{3}</td>
<td>120</td>
<td>3</td>
<td>0.063</td>
</tr>
<tr>
<td>NO\textsubscript{3}</td>
<td>50</td>
<td>5</td>
<td>0.106</td>
</tr>
<tr>
<td>Ca</td>
<td>300</td>
<td>2</td>
<td>0.042</td>
</tr>
<tr>
<td>Mg</td>
<td>30</td>
<td>2</td>
<td>0.042</td>
</tr>
<tr>
<td>Na</td>
<td>200</td>
<td>2</td>
<td>0.042</td>
</tr>
<tr>
<td>K</td>
<td>12</td>
<td>1</td>
<td>0.021</td>
</tr>
<tr>
<td>COD</td>
<td>10</td>
<td>4</td>
<td>0.085</td>
</tr>
<tr>
<td>Zn</td>
<td>3</td>
<td>1</td>
<td>0.021</td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>47</td>
<td>1</td>
</tr>
</tbody>
</table>

EC: Electrical Conductivity; TDS: Total Dissolved Solids; TH: Total Hardness; COD: Chemical Oxygen Demand.
Figure 1. Sampling locations/land uses map (a) and geology map (b) of the study area
The Electrical Conductivity (EC), which depicts the amount of dissolved solids in water, was measured in this study. The EC values of groundwater for drinking purposes (Gw1, Gw3, and Gw4) were lower than the permissible limit of 1000 mho/cm in both seasons. 30 High values of EC (2230 mho/cm) were observed in the sampling points of Gw2, which is in an agricultural land area and near a village. The enrichment of salts in Gw2 with the EC values of 2230 and 2500 mho/cm during pre- and post-monsoon seasons shows that it falls under the medium salt enrichment class (EC: 1500-3000 mho/cm) [37].

The Total Dissolved Solids (TDS) ranged from 230.8 to 300.7 mg/L and 284.8 to 321.8 mg/L in dry and wet seasons, respectively, for the wells with drinking water purposes. These wells fall into freshwater categories (TDS<1000 mg/L) [30]. The measured TDS contents were 1294.9 and 1412.6 mg/L for Gw2 at each season which illustrates the brackish water category (1000-10000 mg/L) [38]. None of the samples taken from Gw1, Gw3, and Gw4 showed TDS values exceeding the permissible limit of 500 mg/L for potable use. Accordingly, all groundwater samples in the dry season exhibited higher TDS values than those obtained from river samples. However, this phenomenon completely reversed in the post-monsoon season so that the TDS amounts in surface water exceeded groundwater levels. Gw1 in the dry season and Gw1, Gw3, and Gw4 in the wet season recorded higher Total Hardness Values (TH) beyond the permissible value of 200 mg/L as CaCO3 and the total alkalinity values were within the safe range of 20-200 mg/L as CaCO3 for potable water [30].

The maximum reported sample of TDS and total alkalinity were from Gw2 with irrigating purposes. TH values were comparatively higher than their corresponding river water samples, indicating the cation-anion exchange through water-rock interaction [36]. Surprisingly, a reverse trend was noticed in the wet season as the amount of EC was lower in groundwater than the detected levels of surface water sampling points. This result may be explained by the fact that the soil is saturated during the wet season. Because of the soil reduction in ion exchange capacity, the concentration of ions decreases, which subsequently ended in a decrease in EC values.

Compared to the dry period, the EC values in both river and groundwater samples are somewhat increased in the wet season, which can be explained by a drop in the river flow resulting from the closing of the upper dam (Latyan dam) gates. Correspondingly, this effect led to a remarkable increase in the concentration of other tested parameters in all sampling points during the wet season (Table 3). The EC values of groundwater for drinking purposes (Gw1, Gw3, and Gw4) lowered the permissible limit of 1000 mho/cm in both seasons. 30 High values of EC (2230 mho/cm) were observed in the sampling point of Gw2, which is in an agricultural land area and near a village. The enrichment of salts in Gw2 with the EC values of 2230 and 2500 mho/cm during pre- and post-monsoon seasons shows that it falls under the medium salt enrichment class (EC: 1500-3000 mho/cm) [37].

The chemical composition of the groundwater and river water samples during pre- and post-monsoon seasons are summarized in Table 3. The pH in river water samples ranged from 8.09 to 8.16 and 7.92 to 8.27 in pre- and post-monsoon seasons, respectively, indicating neutral nature and slight alkalinity of surface water, which were similar to the corresponding pH values of 7.33 to 7.59 and 7.2 to 7.9 observed in groundwater samples recharging from the surface points. However, the pH content of groundwater was slightly lower than their surface values suggesting the dissolution of CO₂ in subsurface aquifers. All the groundwater samples showed the pH values within the permissible limit of 6.5-8.5 30.

The Electrical Conductivity (EC), which depicts the enrichment of dissolved solids in groundwater, varied from 431 to 2230 μmho/cm in the dry season, which was remarkably higher than their corresponding river water samples, indicating the cation-anion exchange through water-rock interaction [36]. Surprisingly, a reverse trend was noticed in the wet season as the amount of EC was lower in groundwater than the detected levels of surface water sampling points. This result may be explained by the fact that the soil is saturated during the wet season. Because of the soil reduction in ion exchange capacity, the concentration of ions decreases, which subsequently ended in a decrease in EC values.
<table>
<thead>
<tr>
<th>Source Type</th>
<th>pH</th>
<th>EC</th>
<th>Turbidity</th>
<th>TSS</th>
<th>TDS</th>
<th>TH</th>
<th>FC</th>
<th>Zn</th>
<th>Cd</th>
<th>Pb</th>
<th>Fe</th>
<th>Cu</th>
<th>Ni</th>
<th>Cr</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rw2</td>
<td>7.9</td>
<td>551</td>
<td>0.17</td>
<td>1.14</td>
<td>0.17</td>
<td>5.3</td>
<td>0.02</td>
<td>11.62</td>
<td>1.16</td>
<td>1.34</td>
<td>4.4</td>
<td>4.4</td>
<td>105</td>
<td>1.16</td>
</tr>
<tr>
<td>Rw3</td>
<td>8.18</td>
<td>333</td>
<td>1.14</td>
<td>0.17</td>
<td>0.17</td>
<td>9.5</td>
<td>0.01</td>
<td>3.62</td>
<td>0.17</td>
<td>4.4</td>
<td>6.1</td>
<td>6.1</td>
<td>105</td>
<td>1.16</td>
</tr>
<tr>
<td>Rw4</td>
<td>8.2</td>
<td>343</td>
<td>0.17</td>
<td>0.17</td>
<td>0.17</td>
<td>14.5</td>
<td>0.02</td>
<td>1.16</td>
<td>4.4</td>
<td>6.1</td>
<td>6.1</td>
<td>6.1</td>
<td>105</td>
<td>1.16</td>
</tr>
<tr>
<td>Gw1</td>
<td>7.59</td>
<td>4112</td>
<td>22.02</td>
<td>0.17</td>
<td>0.17</td>
<td>6.1</td>
<td>0.02</td>
<td>0.34</td>
<td>1.16</td>
<td>4.4</td>
<td>4.4</td>
<td>105</td>
<td>1.16</td>
<td></td>
</tr>
<tr>
<td>Gw2</td>
<td>7.9</td>
<td>4112</td>
<td>22.02</td>
<td>0.17</td>
<td>0.17</td>
<td>6.1</td>
<td>0.02</td>
<td>0.34</td>
<td>1.16</td>
<td>4.4</td>
<td>4.4</td>
<td>105</td>
<td>1.16</td>
<td></td>
</tr>
<tr>
<td>Gw3</td>
<td>8.09</td>
<td>2230</td>
<td>20.81</td>
<td>0.17</td>
<td>0.17</td>
<td>6.1</td>
<td>0.02</td>
<td>0.34</td>
<td>1.16</td>
<td>4.4</td>
<td>4.4</td>
<td>105</td>
<td>1.16</td>
<td></td>
</tr>
<tr>
<td>Gw4</td>
<td>7.2</td>
<td>331</td>
<td>0.17</td>
<td>0.17</td>
<td>0.17</td>
<td>6.1</td>
<td>0.02</td>
<td>0.34</td>
<td>1.16</td>
<td>4.4</td>
<td>4.4</td>
<td>105</td>
<td>1.16</td>
<td></td>
</tr>
</tbody>
</table>

Table 3. River and groundwater quality in the study area.

exists. Overall, we found that the TH and TDS concentrations shifted post-monsoon season in surface or groundwater samples. This result might be due to the leaching of the dissolved materials with monsoon water from the unsaturated zone to the phreatic zone (saturated zone) or activation of dissociation and dissolution process that ultimately enrich waters in ionic contents. Moreover, the influence of lower flow rate during wet seasons on probable higher ions concentrations should also be considered. Ouyang et al. stated that an important parameter contributing to river water quality variation for one season might be insignificant for another season [30]. Bicarbonate was dominant among all the anions, and its concentration slightly rose in the post-monsoon season (Figure 2).

The same trend was observed for other tested anions, as higher amounts were noticed in the wet season. Concentrations of F, Cl, and NO$_3^-$ in groundwater samples were lower than the permissible WHO recommended values for drinking water [30], which indicates the anthropogenic sources are not highly affected the water bodies. However, some authors speculated that nitrate concentration $> 5$ mg/L, as observed in our research, can indi-

![Figure 2](image2.png)

**Figure 2.** Detected anions concentration in groundwater during pre- and post-monsoon season

![Figure 3](image3.png)

**Figure 3.** Detected cations concentration in groundwater samples during pre- and post-monsoon season
Figure 4. Piper diagrams for the chemical composition of groundwater samples

Water type
1. Calcium-rich waters
2. No dominant type
3. Alkali-rich waters
4. Magnesium-rich
5. Bicarbonate waters
6. No dominant type
7. Sulphate waters
8. Chloride waters
9. HCO₃-CO₃ and Ca-Mg
10. SO₄-Cl and Ca-Mg
11. SO₄-Cl and Na-K
12. HCO₃-CO₃ and Na-K
13. Mixing zone

Figure 5. Hydrochemical facies plot illustrating the principal geochemical process in the study area

Water type
1. Ca+Mg > Na+K
2. Na+K > Ca+Mg
3. CO₃+HCO₃ > SO₄+Cl
4. SO₄+Cl > CO₃+HCO₃

cate water contamination by sewage, animal waste, or fertilizers [39, 40]. The maximum concentration of NO$_3^-$ in the study region was found at Gw2 (33.3 and 49.4 mg/L, in dry and wet seasons, respectively) in an agricultural area. These results might be due to the entrance of exogenous contaminants from irrigation water and or fertilizers application over the area. The application of N-fertilizers by farmers in their fields resulted in the leaching of NO$_3^-$ through the soil profile and enriched local aquifers with this ion [41]. Also, SO$_4^{2-}$ content varied from 36.9 to 53.3 mg/L and 52 to 60.2 mg/L in groundwater supplying drinking water, and all were within the permissible range as per by WHO (400 mg/L) [30].

As shown in Table 3, the sulfate concentration in the suppliers’ river water remained relatively stable in various sampling points. Only one sample (Gw2) demonstrated a high concentration of SO$_4^{2-}$ which serves for agricultural irrigation. Higher levels of SO$_4^{2-}$ and Cl$^-$ in Gw2 might be due to the interaction with a saline layer or leakage of irrigation return water [42]; also, its contamination might be related to the proximity to rural areas and contamination with domestic sewage. Sulfate mineral dissolution and sulfide mineral oxidation have been known as the natural sources causing excess SO$_4^{2-}$ in groundwater [43]. Some authors have speculated anthropogenic sources of SO$_4^{2-}$ rather than geogenic origin based on low anhydrite saturation index [44]. In this regard, atmospheric deposition, sulfate-containing fertilizers, bacterial oxidation of sulfur compounds, and other human impacts may result in SO$_4^{2-}$ high values in groundwater [45]. Importantly, all groundwater samples are deficient in minimum fluoride content as recommended by WHO of 0.7 mg/L to prevent dental caries [18]. Therefore, people who use these groundwater sources may be

Figure 6. The ratio of cations [(Na+K)/(Na+K+Ca)] and anions [Cl/(Cl+HCO$_3$)] against TDS
A: Gibbs plots representing the TDS versus Na/(Na+Ca); and B: TDS versus Cl/(Cl+HCO$_3$) in both seasons.
at risk of dental caries. The concentration of HCO$_3^-$ was high in the study area, and all groundwater samples exceeded the WHO standard limits of 120 mg/L for bicarbonate. Appello and Postma speculated that chemical reaction between groundwater and silicate minerals is the main cause of HCO$_3^-$ formation in groundwater. Carbonate-rich rocks, such as crystalline limestone, dolomitic limestone, and calc-granulite, are demonstrated as the main sources for carbonate weathering [46]. Aquifers enrich with high bicarbonate levels during irrigation, rainfall infiltration, and groundwater move through these rocks. The abundance of major anions in groundwater samples was in the order of HCO$_3^-$ >SO$_4^{2-}$ >Cl$^-$ >NO$_3^-$ >F$^-$. Calcium was found as the dominant cation, among others, indicating samples are calcium-rich (Figure 3). Few studies have specifically examined the relation between Ca$^{2+}$ and Mg$^{2+}$ and stated that the calcium content is 3 to 4 fold higher than magnesium in natural waters [48]. In the current study, Ca$^{2+}$ to Mg$^{2+}$ ratio was greater than 3 in most of the examined ground and surface water samples, indicating that they are not polluted. However, this ratio was low for Rw4 that recharges well (Gw2) in both seasons, indicating that the discharges from the local households contaminate this point. Regarding sodium and potassium values, Rw4 and Gw2 were registered with high concentrations. Such elevated levels of these elements do not occur on arable lands [49], which suggests that water composition in these locations may be controlled by wastewater flowing from local households [50].

Calcium and bicarbonate ions were remarkably higher in concentration and the chief principle of elevated levels of EC in the study area. COD values of the groundwater samples ranged from 7 to 16 and <1 to 32 mg/L in pre- and post-monsoon seasons, respectively. Except for Gw1, other groundwater sampling points have registered COD values above the WHO provisional drinking water limit of 10 mg/L in both seasons. It suggests the possible entry of natural organic matter into surface waters that subsequently penetrate aquifers. Also, due to inorganic chemicals in water, a COD test can indicate the amount of oxygen needed to oxidize them [51]. River water demonstrated a relatively higher level of COD as compared with groundwater samples. According to Amneera et al. the elevated COD concentration illustrates polluted water [45]. COD values less than 20 mg/L exhibit unpolluted waters. The higher amounts of COD and Biochemical Oxygen Demand (BOD) were identified in the Jajrood river samples during various seasons in 2009 [52, 53]. The authors concluded that excessive organic contaminants with different sources such as domestic, agricultural, industrial, and naturally influenced water oxygen demand. Based on field observation, excessive COD values (21 and 32 mg/L) in Rw4 during both seasons are related to fertilizers and pesticide application over agricultural lands besides local sewage discharge.

All recorded zinc (Zn) values in water samples were far below the limit set by international guidelines. The Zn values were higher for surface water than their amounts in groundwater samples revealing river water pollution with zinc. The bacteriological analysis (fecal coliform) showed that the groundwater samples were not contaminated (MPN/100 mL <3). Fecal Coliform (FC) values were generally lower than river water samples demonstrating the bacteria screened out by the soil pores and rocks 51 or the bactericidal role of chlorine injected into drinking water wells [54]. Generally, the values obtained in our study are almost similar to those reported by Razmkhah et al. [48], who also emphasized that the morphology of soil, overland runoff, and fertilizers are the main sources of chemical water constituents of the Jajrood river.

**Groundwater quality assessment**

The water quality index has been applied to assess the quality of any water body. To calculate WQI values at each groundwater sampling point, the weight values were assigned for each water quality parameter according to their relative importance in the overall quality of water for drinking purposes. The relative weight of each parameter in the WQI calculation and their corresponding standard levels are shown in Table 2. The parameters of TDS, F$^-$, and NO$_3^-$ were assigned the weight of 5; pH, EC, SO$_4^{2-}$, and COD were assigned 4, taking into account their health significance. K$^+$ and Zn$^{2+}$ were assigned the minimum relative weight due to their least importance in water quality and health-related impacts. The computed WQI of groundwater samples for drinking
Figure 7. Chadha’s plot to reveal the driving geochemical process in the study area

Figure 8. XRD pattern of the soil sample collected from 12m depth of study area
use ranged from 47 to 49.36 and 52.07 to 64.69 in the pre- and post-monsoon seasons, respectively (Table 4). Higher WQI values were observed in post-monsoon seasons as the water quality declined from “excellent” in the pre-monsoon to “good” in the post-monsoon season. Elevated values of WQI in the wet period were related to COD, HCO$_3^-$, TDS, and TH, which all depicted higher Standard Index (SI) values. The amount and composition of dissolved elements in water are of great importance for agricultural use.

Excessive concentrations of dissolved ions in farming water change soil configuration, permeability, and aeration, consequently hardening plant development [55, 56]. The parameters such as sodium percentage (Na%), sodium adsorption ratio (SAR), and residual sodium carbonate (RSC) are essential to determine the suitability of groundwater for agricultural use [57, 58]. Since one of the wells (Gw2) supplies water for agricultural irrigation, and the mentioned indicators were employed to check its water suitability. RSC is used to evaluate the harmful effects of carbonate and bicarbonates on water quality for agricultural purposes [33]. Based on the calculated RSC values (-5.13 and 1.92 meq/L), the well water falls into “safe” and “marginally suitable” categories in pre- and post-monsoon seasons, respectively. The negative values of RSC implied the lower amounts of CO$_2^-$ and HCO$_3^-$ than Ca$^{2+}$ and Mg$^{2+}$ levels in the groundwater. This fact may explain the absence of residual carbonate to react with Na$^+$ to worsen soil quality due to alkali hazards [56, 59]. Considering sodium, the SAR index has been applied for evaluating the suitability of irrigation water [58].

The calculated Na% (41% and 40%) shows that the well water was in “permissible” and “good” conditions for the same periods, respectively. The well water exhibited safe quality based on computed SAR (3.54 and 3.56) in both seasons. Overall, the estimated indexes suggest that the groundwater quality (Gw2) in the study area is suitable for agricultural purposes.

Hydrogeochemical facies (piper)

The Piper diagram was used to investigate the hydrogeochemical facies of groundwater in the Jajrood river basin. As shown in Figure 4, the water of 3 wells (Gw1, Gw3, and Gw4) fall in the zone of calcium-rich and no dominant cation type in pre- and post-monsoon seasons, respectively. Moreover, analysis of the piper diagram revealed that bicarbonate is the dominant ion and the major water type of the study area is the Ca$^{2+}$/Mg$^{2+}$-HCO$_3^-$ (Figures 4 and 5). This finding suggests the presence of temporary hardness in the groundwater of the Jajrood basin. Figure 5 explains the superiority of alkaline earth exceeds alkali (Ca$^{2+}$ + Mg$^{2+}$ > Na$^+$ + K$^+$) and weak acidic anions dominant over strong acidic anions (HCO$_3^-$ > Cl$^-$ + SO$_4^{2-}$). It is also shown in Figure 4 that all groundwater samples except Gw2 in the pre-monsoon period are in zone 9, suggesting temporary hardness is controlling the water quality. Gw2, which is in zone 9, belongs to the mixing zone, where water type can be identified as neither anion nor cation dominant. None of the samples fall under zone 10, indicating non-carbonate hardness belonging to Ca, Mg, SO$_4^{2-}$, Cl does not exist. Furthermore, no samples fell under zone 11 and 12, suggesting that halite dissolution (saline) or alkali carbonate enrichment were not affecting water type [57]. According to Mechal

Table 4. Classification of groundwater based on calculated Water Quality Index (WQI)

<table>
<thead>
<tr>
<th>Sample No.</th>
<th>WQI (Water Type) Pre-monsoon</th>
<th>Post-monsoon</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gw1</td>
<td>49.36 (Excellent water)</td>
<td>52.07 (Good water)</td>
</tr>
<tr>
<td>Gw3</td>
<td>47.00 (Excellent water)</td>
<td>60.95 (Good water)</td>
</tr>
<tr>
<td>Gw4</td>
<td>49.32 (Excellent water)</td>
<td>64.69 (Good water)</td>
</tr>
</tbody>
</table>

Table 5. Inductively Coupled Plasma (ICP) results of elements content in the investigated soil

<table>
<thead>
<tr>
<th>Type of Elements</th>
<th>Na</th>
<th>Mg</th>
<th>Al</th>
<th>Si</th>
<th>P</th>
<th>S</th>
<th>Cl</th>
<th>K</th>
<th>Ca</th>
<th>Hg</th>
</tr>
</thead>
<tbody>
<tr>
<td>%</td>
<td>0.60</td>
<td>1.25</td>
<td>4.00</td>
<td>17.50</td>
<td>0.07</td>
<td>0.01</td>
<td>0.01</td>
<td>1.40</td>
<td>10.58</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Type of Elements</th>
<th>Fe</th>
<th>Co</th>
<th>Ni</th>
<th>Cu</th>
<th>Zn</th>
<th>As</th>
<th>Se</th>
<th>Cr</th>
<th>Mn</th>
<th>Cd</th>
</tr>
</thead>
<tbody>
<tr>
<td>%</td>
<td>2.00</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>0.01</td>
<td>0.10</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>
et al. as the CO₂ hydrolyzed soil silicate, HCO₃⁻ releases in the groundwater. In this research, the dominance of Ca²⁺-HCO₃⁻ water type illustrates the possible hydrolysis of silicate minerals by CO₂ [56]. Katz et al. also stated that the Ca²⁺/Mg²⁺ ratio greater than 2 indicates that the hydrolysis of silicate minerals is the chief reason for high Ca²⁺ and Mg²⁺ amounts in the groundwater [57]. According to Table 1, all samples have Ca²⁺/Mg²⁺ ratios greater than 2 revealing the hydrolysis of silicate minerals could lead to Ca²⁺/Mg²⁺ release into groundwater.

The Mechanism controlling groundwater chemistry (chadha and gibbs)

Gibbs proposed the geochemistry of groundwater with respect to atmospheric precipitation, rock dissociation, and evaporation. However, Li et al. reported the downside of the Gibbs diagram. They argued that the impacts of human activities on groundwater chemistry have been ignored [59].

This diagram represents the ratio of cations [(Na+K)/(Na+K+Ca)] and anions [Cl/(Cl+HCO₃⁻)] against TDS. Figure 6 specifies that most of the water samples during both periods belonged to rock dominance, with the TDS values below 1000 mg/L, affecting groundwater chemical constitution. No samples were classified as precipitation or evaporation dominant, indicating the limited role of atmosphere and the dissolution of evaporates like halite (NaCl), gypsum (CaSO₄·2H₂O), and mirabilite (Na₂SO₄) in the input of solutes [60, 61]. The post-monsoon season did not alter the hydrochemical facies in the study area. Since high TDS content was accompanied by a notable increase in HCO₃⁻ concentration, the role of evaporation is negligible. The geology and environment of water affect the type and amounts of salts in water bodies [62]. Chadha proposed a hydrochemical diagram reflecting various hydrochemical processes involved in water quality [12]. The differences between alkaline earth (Ca²⁺+Mg²⁺) and alkaline metals (Na⁺+K⁺) in meq/L for each sampling point were plotted against the difference between weak acidic anions (HCO₃⁻+CO₃²⁻) and strong acidic anions (Cl⁻+SO₄²⁻). Then, the result is depicted in Figure 7. Chadha suggested four processes controlling the quality of water: base ion exchange water, reverse ion exchange water, recharging water, and seawater types [12]. The position of data points in Figure 7 demonstrates that recharging water was the driving process of the chemical composition of water in both seasons. Recharging waters are shaped when water penetrates the soil and dissolves carbonate and calcium in the subsurface layers. However, based ion exchange was the dominant process for one sampling point (Gw2) in the wet period, which represents the base ion exchange reaction path of groundwater. In this case, Ca²⁺-HCO₃⁻ type water mixes with Na⁺-Cl⁻ water type, which finally produced Na⁺-HCO₃⁻ type. The high values of Na⁺ and Cl⁻ for site Gw2 in post-monsoon (Table 1) also confirm this phenomenon. The high values of chloride ions in the water represent the influence of agrochemicals and domestic wastewaters [62]. Given that the study area is in agricultural communities, applying organic and inorganic fertilizers might pollute the groundwater by this ion.

Quality in relation to subsurface soil structure and composition

Natural mineral weathering dissolution can be the main reason for high-level EC related to dissolved solids in groundwater bodies. In this regard, some subsurface soil samples were analyzed using XRD, XRF, and ICP for the presence of the detailed minerals/metals, their concentrations, and the soil composition. According to Figure 8, the main phases of the soil were quartz, calcium carbonate, and dolomite. Also, calcium aluminum silicate and illite constituted the minor phases. The high concentrations of Ca²⁺ and Mg²⁺ detected in groundwater samples compared to their surface values, especially during the pre-monsoon period, can be explained by the soil enrichment of these cations (Table 5). Besides, XRD data confirms the presence of calcium carbonate, which forms temporary hardness as earlier presented by the results of the Piper diagrams. The result of the XRF analysis is depicted in Table 6. As shown, Si, Ca, Al, Fe, and Mg contributed to the highest percentage of soil composition among other identified elements, confirming their geogenic origins over anthropogenic sources. Other studies have shown that the high content of Fe in soils can

<table>
<thead>
<tr>
<th>Type of Elements</th>
<th>Ba</th>
<th>Ca</th>
<th>Co</th>
<th>Cr</th>
<th>Cu</th>
<th>Fe</th>
<th>K</th>
<th>Mg</th>
<th>Mn</th>
<th>Ni</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concentration (mg/kg)</td>
<td>260</td>
<td>102100</td>
<td>7</td>
<td>20</td>
<td>18</td>
<td>15200</td>
<td>19000</td>
<td>11200</td>
<td>885</td>
<td>20</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Type of Elements</th>
<th>Pb</th>
<th>P</th>
<th>Se</th>
<th>Zn</th>
<th>Ag</th>
<th>As</th>
<th>Bi</th>
<th>Cd</th>
<th>Ga</th>
<th>Hg</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concentration (mg/kg)</td>
<td>2.00</td>
<td>510</td>
<td>&lt;0.1</td>
<td>&lt;0.01</td>
<td>0.01</td>
<td>&lt;0.01</td>
<td>&lt;0.01</td>
<td>0.01</td>
<td>0.10</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

Table 6. X-ray Fluorescence Spectrometry (XRF) analysis of the collected soil samples
contribute to mutagenic/carcinogenic effects on the health of the human being [61]. Because Fe and Al occur naturally at plentiful levels; thus, they hardly originated from anthropogenic inputs. Furthermore, according to Table 5, most detected heavy metals revealed the <0.01% values in the analyzed soil, expressing no pollution. Similarly, the ICP-OES data of sampled soil (Table 5) showed the abundance of Ca+2, K+2, Fe+2, and Mg+2 concentrations, which are in accordance with their high observed contents in XRF analysis. Since no heavy metal was identified as soil constituents, their presence could be attributed to anthropogenic sources such as fertilizers, sewage, and industrial discharge to the river. Although the concentration of most heavy metals was low in the sampled soil, Zn, Ni, and Cr amounts were noticeable, suggesting the soil of the study area is affected by anthropogenic activities. The lower detected concentration of zinc in groundwater compared to the river water samples indicate the role of the soil in the low amount of this metal and its retention in the soil matrix. An elevated zinc level has been found in groundwater in regions with mining activities, attributed to industrial waste disposal and the introduction of sludge to the environment [59].

4. Conclusions

In this research, various hydrogeochemical techniques and quality indices are applied to define the suitability of groundwater quality for drinking and irrigation purposes. While preliminary findings suggest that natural processes control the geochemistry of the study region, in some locations revealed anthropogenic sources were also involved. The groundwater of the study region is neutral to slightly alkaline due to the intrusion of HCO3 ions in the groundwater aquifer, as most of the samples exhibited higher levels of bicarbonate. This result is further supported by the subsurface XRD soil analysis in which calcium carbonate (Ca3C6O18) was recognized as one of the main soil components. Hydrogeochemistry analysis reveal that the order of ions abundance were Ca+2 > Na+ > Mg+2 > K+ as cations and HCO3 > SO42- > Cl- > NO3 > F- as anions. Except for HCO3 and COD, all water wells are fit with little precaution for drinking purposes, and the rest of the studied parameters are within the permissible limits set by WHO. Water quality indices and ratios for drinking and agricultural purposes demonstrated the safety of all samples for their determined uses. Contamination with a low nitrate concentration exhibits the input of anthropogenic activities like domestic sewage, animal excreta, and fertilizers application. The results from the Piper trilinear diagram demonstrated that nearly all samples belonged to Ca-Mg-HCO3 water type, which forms temporary hardness.

Ethical Considerations

Compliance with ethical guidelines

There were no ethical considerations to be considered in this research.

Funding

This research did not receive any grant from funding agencies in the public, commercial, or non-profit sectors.

Authors' contributions

All authors equally contributed to preparing this article.

Conflict of interest

The authors declared no conflict of interest.

Acknowledgments

The authors would like to thank for the support of the Water and Wastewater Reference Laboratory affiliated to Iran University of Medical Sciences for conducting water samples analysis and special thanks to the Ministry of Energy and Rural Water Supply Company for providing the required information.

References


